Two cross section fitting procedures may be used. In one, fitted cross sections are provided for each isotope which are independent of position but may be changed at any time in life. Thermal self-shieding factors can be specified for each isotope taking on different values for various subregions. A correction is made to the epithermal absorption and removal cross sections to account for the U²³⁸ (Hf¹⁷⁷) resonances. A second procedure allows the fast cross sections of U235 and the isotopes in the U²³⁸ (Hf¹⁷⁷) chain to be fitted as the inverse of a bilinear function in the isotopic densities of U²³⁵ and U²³⁸ (Hf¹⁷⁷). Assuming that the "mixed number density" scheme is used in the thermal group, the cross sections of U²³⁸ (Hf¹⁷⁷), Xe¹³⁵, and Sm¹⁴⁹ are fitted as linear functions of the density of U²³⁵. In either of the schemes, explicit treatment of B¹⁰ plates is included by use of fictitious diffusion and absorption cross sections fitted as polynomials in the B¹⁰ density. Variable selfshielding factors are allowed for U²³⁵ and B¹⁰ fitted as the inverse of a linear function in the fraction of the isotope remaining at the various points in the mesh.

Maximum or maneuvering xenon calculations may be performed at various times in the calculation.

Provision is included for the study of fuel replacement. The xy version is also designed to provide the fluxweighted macroscopic data required to do the flux synthesis by the ZIP program.

- 6. Restrictions on the complexity of the problem: No more than 178 mesh subdivisions may be used along the horizontal axis or 201 mesh divisions along the vertical axis with a limit of 20,000 on the product of the two. A maximum of 150 material compositions or subregions of interest can be defined and 2, 3, or 4 lethargy groups can be used. Input is compatible with the other diffusion theory programs on the Philco-2000 computer such as WANDA-5, PDQ-4, CANDLE, etc. The program is intended to operate within the BKS System.
- 7. Typical running time: The typical running time for one time step is greater than that of the corresponding PDQ-4 problem by a factor of one-third.
- 8. Present status: In use
- References: O. J. Marlowe, Nuclear reactor depletion programs for the Philco-2000 computer. WAPD-TM-221 (January 1961).

R. B. Smith and C. H. Hunter, The BKS system for the Philco-2000 computer. WAPD-TM-233 (April 1961).

W. R. Cadwell and H. P. Henderson, Input preparation for diffusion-depletion programs on the Philco-2000 computer. WAPD-TM-238 (January 1961).

W. R. Cadwell, PDQ-4—A program for the solution of the neutron-diffusion equations in two dimensions on the Philco-2000. WAPD-TM-230 (June 1961).

C. J. Pfeifer and F. R. Urbanus, ZIP-2-A one-dimensional few-group synthesis nuclear reactor depletion program for the Philco-2000 computer. WAPD-TM-228 (November 1961).

10. Material available from Philco:

Binary program deck

Symbolic program tape Referenced documents W. R. CADWELL, O. J. MARLOWE L. M. CULPEPPER, J. P. DORSEY, J. T. MANDEL, H. H. MITCHELL, C. J. PFEIFER, M. C. SUGGS Westinghouse Electric Corporation Bettis Atomic Power Laboratory Pittsburgh, Pennsylvania

WANDA-5

- 1. Name of program: WANDA-5
- 2. Computer for which program is designed: Philco-2000 Programming system: TAC
- 3. Nature of problem solved: Numerical solution of the one-dimensional few-group neutron diffusion equations. One to eight energy groups may be used and rectangular, cylindrical, or spherical geometry formulations are available. The flux or its derivative may be set to zero at the boundaries. The program will vary buckling or poison cross section in any subset of regions, or the position of one or more interfaces separating regions to find a specified critical eigenvalue. One iteration or fixed source problems may be calculated and adjoint solutions may be obtained. All one-group problems are treated as one iteration problems.
- 4. Method of solution: The diffusion equations are replaced by difference equations giving rise to a tridiagonal matrix inverted by Gauss elimination. Extrapolation procedures are incorporated in the source calculation to accelerate convergence.
- 5. Restrictions on the complexity of the problem: Limited to 500 points, 100 regions, eight groups, and 2000 grouppoints. This program is intended to operate within the BKS system.
- 6. Typical running time: 1 min
- 7. Present status: In use
- References: O. J. Marlowe and M. C. Suggs, WANDA-5— A one-dimensional neutron diffusion equation program for the Philco-2000 computer. WAPD-TM-241 (November 1960).
 W. R. Cadwell and H. P. Henderson, Input preparation for diffusion-depletion programs

on the Philco-2000 computer. WAPD-TM-238 (January 1961). R. B. Smith and C. H. Hunter, The BKS sys-

tem for the Philco-2000 computer. WAPD-TM-233 (April 1961).

9. Material available from Philco:

Binary program deck

Symbolic program tape

Referenced documents

O. J. MARLOWE

Westinghouse Electric Corporation Bettis Atomic Power Laboratory Pittsburgh, Pennsylvania

ZIP-2 and ZIP-3

- 1. Name of programs: ZIP-2, ZIP-3
- 2. Computer for which programs are designed: Philco-2000 Programming system: TAC
- 3. Nature of problem solved: ZIP-2 and ZIP-3 are diffusion theory synthesis depletion programs for design calculations. The synthesis technique is a method of approximating a solution to the three-dimensional, few-group diffusion theory equations by "stacking" a